subsequent direct sequence assays were performed from each purified DNA extract. The mtDNA sequences derived from each bone were therefore multiply verified through independent samples, extractions, amplifications and sequence determinations. In all cases for each bone in the panel, the replicated mtDNA sequences were consistent across all samples and extractions. The veracity and integrity of the aurochs sequence haplogroup is strongly supported by the congruity of the bones analysed in Dublin (CHWF, TP65, CPC98 and NORF) with those previously examined in Oxford (D740 and D812).

Analysis of mitochondrial sequence data

We aligned sequences by eye. Data from previous studies^{5-7,17} were also included and novel sequences were submitted to GenBank (accession numbers AF336383-AF3366748). We constructed a neighbour-joining phylogeny using uncorrected distances with gapped positions excluded and the Neighbour program in the Phylip package¹⁹. Arlequin 2.0 (ref. 20) was used to estimate nucleotide diversity values for each population, to estimate parameters and goodness of fit from mismatch distributions and to compute Fu's Fs test of selective neutrality¹⁰. Phylogenetic analysis was also performed using reduced median networks, constructed manually according to ref. 21. Sites prone to hypermutablity were identified using a network of zebu and taurine samples examined in ref. 6. For the reduced median networks shown, most of the reductions (71%) were at these sites. The mutation rate for the 240-bp region of the cattle D-loop was calculated using the estimate of the transition/transversion ratio for the sequence data presented here (61/1). This agrees well with previous estimates on smaller data sets 57/1 (ref. 6) and 41/1 (ref. 7), and as the new rate incorporates data presented in those papers it is used in preference to either. The Bison-Bos divergence is considered to be around 1 Myr. The three transversions observed in the 240-bp region between the Bison-Bos groups constitute the equivalent of 183 transitions. The one-lineage rate was estimated as 38% per Myr or one substitution per 10,928 yr. The central 95% credible region for the expansion time was calculated for each of the four main cattle haplogroups using the program CRED²².

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The highly reduced genome of an enslaved algal nucleus

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Chromophyte algae differ fundamentally from plants in possessing chloroplasts that contain chlorophyll c and that have a more complex bounding-membrane topology¹. Although chromophytes are known to be evolutionary chimaeras of a red alga and a non-photosynthetic host¹, which gave rise to their exceptional membrane complexity, their cell biology is poorly understood. Cryptomonads are the only chromophytes that still retain the enslaved red algal nucleus as a minute nucleomorph²⁻⁴. Here we report complete sequences for all three nucleomorph chromosomes from the cryptomonad Guillardia theta. This tiny 551kilobase eukaryotic genome is the most gene-dense known, with only 17 diminutive spliceosomal introns and 44 overlapping genes. Marked evolutionary compaction hundreds of millions of years ago^{1,4,5} eliminated nearly all the nucleomorph genes for metabolic functions, but left 30 for chloroplast-located proteins. To allow expression of these proteins, nucleomorphs retain hundreds of genetic-housekeeping genes5. Nucleomorph DNA replication and periplastid protein synthesis require the import of many nuclear gene products across endoplasmic reticulum and periplastid membranes. The chromosomes have centromeres, but possibly only one loop domain, offering a means for studying eukaryotic chromosome replication, segregation and evolution.

Soon after the symbiogenetic origin of chloroplasts from cyanobacteria¹ to form the common ancestor of green plants, red and glaucophyte algae (kingdom Plantae^{6,7}), even more complex eukaryotic cells arose by secondary symbiogenesis^{1,3,4} (Fig. 1). Such chimaeric integration of two evolutionarily distant eukaryotic cells occurred independently in the common ancestor of cryptomonads and other chromophytes, in which the endosymbiont was a red alga, and in chlorarachneans, which acquired a green alga^{1,3,4}. In both cryptomonads and chlorarachneans, a flagellate host contributed the nucleus, endomembranes and mitochondria to the chimaera, whereas the photosynthetic endosymbiont provided its chloroplast, plasma membrane (the periplastid membrane^{1,3,4}) and a second nucleus (the nucleomorph), which became miniaturized³⁻⁵. The nucleomorph of both groups kept its envelope, nuclear pores⁸ and three minute chromosomes⁹. In the ancestor of cryptomonads and chromobiotes (treated as kingdom Chromista^{6,8}) but not chlorarachneans, the former food vacuole membrane originally enclosing the enslaved endosymbiont apparently fused with the nuclear envelope10, placing it in the rough endoplasmic reticulum8,10 (RER; Fig. 1). Cryptomonad cells depend on four genomes, each encoding distinct protein synthesis machineries in discrete

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letters to nature

compartments, between which proteins are translocated. Until now, understanding how these genomes cooperate has been limited by the availability of only partial sequences for nucleomorph genomes^{11,12}.

We report the first complete genome sequence of a nucleomorph (551,264 base pairs (bp)), which proves conclusively that it is a vestigial nucleus¹³ and that cryptomonads comprise one eukaryotic cell nested in another. Gene density is extremely high (1 gene per 977 bp) and non-coding regions are ultrashort, with only one pseudogene φ *rpl24* (Fig. 2). The six chromosome ends are identical repeats, comprising telomeres ([AG]₇AAG₆A)₁₁, 5S and 28S/5.8S/ 18S ribosomal RNA genes and five open reading frames (ORFs); the ubiquitin conjugation enzyme gene is repeated at five ends, and the TATA-box binding protein gene (*tfIId*) at three ends. Except in these repeats and five central 200-500-bp regions (possibly centromeric), intergenic spacers have few, if any, nucleotides. Forty-four genes overlap by up to seventy-six nucleotides (Fig. 2). One gene has three copies, one on chromosome 1 (ORF 160a) and two on chromosome 3 (ORF 160b,c); the only other repeated genes are in the termini. Coding regions of some genes are shorter than their homologues in other organisms.

Only 17 protein-coding genes contain spliceosomal introns (42-52 bp long; see Supplementary Information), all located in the 5' region, as in yeast¹⁴, many immediately after the initiator AUG. Eleven are in ribosomal protein genes, as in yeast where their splicing negatively regulates messenger RNA levels¹⁴. Like the even shorter pygmy introns of chlorarachnean nucleomorphs¹², they have standard GT/AG boundaries. Twelve transfer RNA genes have protein-spliced introns. The marked contrast between the effective elimination of non-coding DNA from cryptomonad nucleomorphs and the accumulation of vast amounts of noncoding DNA in coexisting cryptomonad nuclei indicates that nuclear non-coding DNA in general is functional and positively selected⁵, and is not purely selfish or junk. Chromosomal A+T content varies, suggesting that there have been three different mutational and selective pressures on base composition: on the terminal repeats (45.5% G+C); on housekeeping genes with a very low G+C content (23%); and on transfer RNA genes and genes for plastid proteins with intermediate G+C content (35%).

The function of 219 of the 464 putative protein-coding genes is unknown, but 31 have convincing database matches—11 to cyanobacteria and 20 to eukaryotes. Retention of the latter, even in this exceptionally compacted genome, shows that they must have important functions in all eukaryotes. The other 245 genes have homologues of known function, mostly for chromosome reproduction or gene expression, with very few for cytoplasmic functions such as protein assembly and degradation, signal transduction/ regulation, cell-cycle control and membrane transport (Fig. 3). Only one gene for metabolism (carotenoid synthesis) was found. There are 47 different genes for non-mRNAs (rRNA, tRNA, small nuclear RNA, small nucleolar RNA).

Three conclusions can be drawn about intergenome cooperation in these complex eukaryote/eukaryote chimaeras. First, most identified genes (> 250/302) are needed simply for self-perpetuation of the nucleomorph and its periplastid ribosomes. End products directly useful to the rest of the cell are encoded by only a few genes including 30 chloroplast proteins, 3 transporters (Sut1 for sulphate; Kea1 for potassium; Rli1, an ABC transporter), an anabolic enzyme (Ggt), and a few regulatory enzymes. Second, even fewer cryptomonad plastid proteins are encoded by the nucleomorph than by the plastid genome¹⁵, so at least a thousand more¹ must be imported into the plastid across four membranes, not two membranes as for nucleomorph-encoded ones. Third, as certain well-conserved genes essential for nucleomorph functions are absent from the nucleomorph genome, these functions must be provided by nuclear genes and imported into the periplastid compartment.

It was not known previously that nuclear gene products not destined for the chloroplast were imported into the periplastid space⁴. As in mitochondria and chloroplasts, DNA polymerase genes are absent. DNA polymerases must be nuclear encoded and imported across the RER and periplastid membranes, and onwards into the nucleomorphs through their nuclear pores. As we identified so few genes for transporters, most must be encoded by nuclear genes or nucleomorph ORFs. As chromobiotes lost the nucleomorph, they are likely to have homologous transport proteins in their nuclei. If chromists are sisters of alveolate protozoa, sharing a photosynthetic common ancestor^{1,4,16}, some may be also present in Sporozoa, such as malaria parasites, and important for their periplastid membranes and as potential drug targets.

Under 10% of the genes encode end-product functions⁵ that are useful to the rest of the cell. Originally three end-product functions



Figure 1 Secondary symbiogenetic origin and membrane topology of cryptomonads. After the primary endosymbiotic incorporation of a cyanobacterium to form the first chloroplast (green), many of its genes were transferred into the host nucleus. After this ancestral plant diversified to form green plants, red algae and glaucophytes, more complex algae were formed by independent secondary symbioses involving green or red algae, after which many or all plastid protein genes were transferred from the algal to the secondary host's nucleus. Shown here is the symbiosis of a red alga to form cryptomonads, where (as in all chromists) the food vacuole membrane fused with the RER. This fusion did not occur in alveolates, chlorarachneans or euglenoids. Former cyanobacterial genes now inserted in nucleomorph or nuclear chromosomes are shown in green, and former red algal genes now in the host nucleus in red. In cryptomonads, the chloroplast and nucleomorph (former red algal nucleus) are topologically in the periplastid space (starch- and ribosomecontaining residual cytoplasm of the former red algal cell, vellow) in the periplastid membrane (former red algal plasma membrane), which is located in the lumen of the host's rough endoplasmic reticulum (RER). Chloroplast proteins are coded by three genomes (chloroplast, nucleomorph and nucleus) and mitochondrial proteins by two genomes (mitochondrion and nucleus). Nucleomorph and periplastid proteins are coded by two genomes (nucleus and nucleomorph). Coloured dots indicate protein translocation pathways in the periplastid complex: nuclear- or nucleomorph-encoded proteins targeted to the chloroplast are green; nuclear-encoded proteins imported into the periplastid space, and both nuclear- and nucleomorph-encoded proteins imported into the nucleomorph, are red. Mitochondrial proteins (brown) are encoded by both nuclear and mitochondrial genomes.



Figure 2 The cryptomonad nucleomorph chromosomes showing gene locations. Colours show the functional categories of the genes specified in Fig. 3. The number of overlapping nucleotides between adjacent genes is indicated beside the brackets. Each chromosome

is displayed as if broken into two fragments near its midpoint. Labels on the left of each chromosome indicate genes transcribed towards its left end (top left); labels on the right show those transcribed towards the right end (bottom right).

letters to nature

were envisaged: plastid proteins not made in the chloroplast itself or in the host cytosol; periplastid space functions in starch metabolism; and periplastid membrane transporters mediating metabolite exchange between host and symbiont. We identified 30 genes encoding proteins for chloroplast function, including two known proteins¹¹, but very few for periplastid functions.

Sequence comparisons show that two nucleomorph genes for plastid proteins are eukaryotic inventions (Iap100, Met), one (CbbX) is α -proteobacterial¹⁷, and the others are cyanobacterial. Only two encode molecules directly needed for photosynthesis: an electron transfer molecule (rubredoxin)¹¹ and a carotene-binding protein (Hlip). Some genes are needed for chloroplast division (FtsZ¹¹) or gene expression (ribosomal protein Rps15, products for DNA and RNA metabolism, and a factor for RuBisCo expression, CbbX). Others encode components of the chloroplast proteinimport machinery (Iap100, Tic22), translocation into the thylakoid lumen (Tha4, SecE), chaperonins (Cpn60, Hcf136) and a protease (ClpP).

All are essential for chloroplast function, explaining why the nucleomorph has persisted for hundreds of millions of years. Compared with their cyanobacterial homologues, the nucleomorph-encoded plastid proteins have amino-terminal extensions (transit peptides; see Supplementary Information) to specify import into the chloroplast, although only those of Hcf136, Cpn60, Tic22 and GyrB were recognized by the search tool ChloroP (http://www.cbs.dtu.dk/services/ChloroP/). Nevertheless, *in vitro*



Figure 3 The fraction of genes in each functional category and their classification as endproduct or genetic housekeeping genes. The genes in each category are listed in the Supplementary Information.

the N-terminal extension of rubredoxin acts as a transit peptide with pea or cryptomonad plastids and is removed after import¹⁸.

Acquiring solar-powered carbohydrate synthesis is the main advantage of enslaving a photosynthetic cell. As glucose produced by degrading periplastid starch is used mainly in the host cytosol, starch metabolism must be responsive to its needs. If the phosphatidylinositol-4-OH kinase (PI(4)K) is associated with the periplastid membrane, it may mediate such signalling through the second messenger inositol trisphosphate. In response, nucleomorph-encoded GTP-binding proteins (GbIp, GTPbp) may regulate periplastid starch metabolism and protein synthesis, as may a protein kinase similar to glycogen synthase kinase; this AMPactivated kinase (AakB) might control starch degradation (like its homologue in animal glycogen degradation). The nucleomorph has four other kinase genes (gs, mps1, snf1, snf2). Protein phosphatase, Pp1, may function complementarily in starch regulation and/or kinase-regulated cell-cycle controls. The presence of a geranylgeranyl-transferase (Ggt) lacking a transit peptide indicates that at least one intermediate stage in carotenoid synthesis may occur in the periplastid space. As plastids typically make carotenoids using isoprenoid precursors from the host, intermediates presumably traverse the periplastid space.

The red algal symbiont lost mitochondria, peroxisomes, lysosomes, Golgi membranes and most metabolic enzymes. Possibly it also lost all ER lumenal proteins and protein glycosylation, perhaps facilitating an unprecedented simplification of eukaryotic membrane-protein targeting. This red algal vestige may differ from true eukaryotic and bacterial cells by lacking co-translational protein insertion into membranes by the signal recognition particle (SRP): the nucleomorph outer membrane, phylogenetically equivalent to RER⁸, uniquely lacks obvious surface ribosomes^{13,19}, and we found no 7S SRP RNA or other genes for SRPs. Perhaps nucleomorph envelope and periplastid membrane proteins are inserted posttranslationally, as in mitochondria and plastids¹ (which analogously both lost the ancestral eubacterial SRP RNA), using the nucleomorph-encoded Hsp70 chaperone.

Genes for DNA polymerase are absent, but there are genes for its sliding clamp (Pcna), a replication co-factor (RfC) for primer extension, and a RecA-like protein (Rad51). Replicon origins might lie in the terminal repeats proximal to the 18S rRNA genes, where there is more non-coding DNA than elsewhere; if there are no others, each chromosome will consist of just two 80–100-kilobase (kb) replicons. As eukaryotic replicons average 100 kb (ref. 20), no origins need lie in unique chromosome regions where intergenic spacers are almost all shorter than the minimal 250 bases needed for origins.

Genes for three core histones (H2b, H3, H4) are present. H2b is

Table 1 Codon usage, introns and chromosomal location of tRNA genes											
Codon	Chr*	%	Codon	Chr*	%	Codon	Chr*	%	Codon	Chr*	%
TTC F	2†	21.4	TCC S		5.6	TAC Y	1†	20.2	TGC C	3†	24.6
TTT F		78.6	TCT S	2†	34.1	ΤΑΤ Υ		79.8	TGT C	2†	75.4
TTA L	3†	50.8	TCA S		34.1	TAA Z		60.0	TGA Z		20.0
TTG L	2†	17.6	TCG S	1	5.6	TAG Z		20.0	TGG W	3†	100
CTC L		1.9	CCC P		12.6	CAC H	2	18.4	CGC R		0.8
CTT L	3	14.9	CCT P	3	32.3	CAT H		81.6	CGT R	3	2.4
CTA L	1	12.2	CCA P		48.3	CAA Q	1	78.2	CGA R	2	8.1
CTG L		2.7	CCG P	1	6.9	CAG Q	2	21.8	CGG R		2.4
ATC I		12.9	ACC T		10.6	AAC N	3	23.6	AGC S	2	6.0
ATT I	1	48.6	ACT T	2	34.1	AAT N		76.4	AGT S		14.5
ATA I	2	38.5	ACA T	3	49.2	AAA K	2	87.1	AGA R	1†	77.4
ATG M _i	3	100	ACG T	3	6.2	AAG K	2	12.9	AGG R	3†	8.9
ATG Me	3†	100									
GTC V		7.5	GCC A		9.1	GAC D	1	18.6	GGC G	3	9.2
GTT V	2†	48.4	GCT A		32.6	GAT D		81.4	GGT G		36.2
GTA V	1	34.4	GCA A	3	50.8	GAA E		84.1	GGA G	3	48.5
GTG V		9.7	GCG A		7.6	GAG E		15.9	GGG G		6.1

 $M_{\text{e}},$ elongator methionine; $M_{\text{i}},$ initiatior methionine

* Chromosome on which tRNA gene is located.

tRNA gene contains non-spliceosomal intron.

exceptionally divergent with its normally acetylated basic Nterminal tail mostly deleted, so nucleosomes may be simplified compared with those of typical nuclei. Histones probably undergo acetylation, as genes for a histone acetyltransferase (Hat) and deacetylase (Hda) are found, and all H3 and H4 acetylation sites are conserved. H3 and H4 genes are adjacent, divergently transcribed and relatively well-conserved, but have evolved more rapidly than nuclear homologues (like most nucleomorph genes). Although H2a and H1 are undetectable, neither is likely to be dispensable, as a string of nucleosomes, unlike a 30-nm filament requiring H1, would be too long for segregation without them.

One gene each for α -, β -, and γ -tubulin is present¹¹, and in addition to γ -tubulin three other centrosomal proteins²¹ are nucleomorph-encoded—Ranbpm¹¹; Hsp82 (in the cytosolic Hsp90 family) and Hsp 70 (cytosolic family). As centrosome activation involves the cyclinB–cdc2 complex²² (both nucleomorph encoded), mitotic control is present. Nuclear localization signals on Hsp82 and Hsp70 suggest that the centrosome is inside the nucleomorph, consistent with the intact envelope during division¹⁹ and the intranuclear red algal spindles. Presence of a centromere-specific H3-like histone (Cenp-A) suggests that nucleomorph have a relict mitotic apparatus¹¹ despite the lack of ultrastructural evidence for a spindle¹⁹.

We cannot be certain of centromere positions. Only the terminal repeats have significant non-coding DNA regions repeated on all chromosomes, but we doubt that they are centromeric because two per chromosome would upset segregation. Assuming nucleomorph centromere sizes to be similar to *Saccharomyces cerevisiae*²³, chromosome 1 has two non-genic regions long enough to be centromeric (520 nucleotides between ORFs 147 and 471; 540 nucleotides between ORFs 244 and 446); chromosome 2 has one (~200 nucleotides between ORFs 80a and 231); chromosome 3 has two (374 nucleotides between ORFs 141 and 180; ~300 nucleotides between ORF 62 and Ebi). As these candidate centromere regions are near chromosome midpoints and other intergenic regions seem too short, all three chromosomes may be metacentric.

Eukaryotic chromosomes are linear arrays of looped domains attached to a nuclear skeleton by their replicon origins²⁴. Each nucleomorph chromosome may constitute a single loop domain, being similar in size. If nucleomorph histones compact the DNA into typical 30-nm chromatin threads, the longest chromosome would be 1.5 μ m in the absence of higher-order condensation; if metacentric, each chromosome arm would be 0.75 μ m. As nucleomorphs are about 1.5- μ m across and longer when dividing¹⁹, extra folding would be unneccessary for mitotic segregation, unlike for other eukaryotic chromosomes. Compacted chromosomes cannot be seen during nucleomorph division¹⁹. Analogous folding constraints may explain why cryptomonad and chlorarachnean nucleomorphs retained three short chromosomes^{3,9,12} instead of aggregating them into a larger one, which would be feasible only with higher-order mitotic compaction.

Genes for key cell-cycle control proteins include a replicationlicensing protein (Mcm2), which mediates the G1/S-phase transition²², and a cyclin-dependent Cdc2 kinase and its cyclin B, which are involved in the G2/M-phase checkpoint. The persistence of a nucleomorph-specific cell-cycle kinase system, even in this simplified cell vestige, emphasizes that it remains conceptually equivalent to a cell, despite its long integration and loss of metabolism and most genes. The nucleomorph encodes ubiquitin (fused to two ribosomal proteins), the ubiquitin-fusion-degradation enzyme (Ufd), and three E2 enzyme subunits (Ubc2, Ubc4, UbcE2). The cell-cycle regulatory role of proteasomes (degrading cyclin B to ensure exit from mitosis, and also controlling exit from S phase²⁵) may be the key reason why such complex multiprotein assemblies as proteasomes and their associated ubiquitin pathway are conserved in the periplastid space, otherwise so reduced in normal cytosolic functions. Supporting this is a gene for an AAA-ATPase (Cdc48; plus a partially related gene *cdc48a*) implicated in ubiquitindependent mitotic events, including membrane fusion²⁶. There are 21 nucleomorph genes for different subunits of the 20S core proteasome and its 19S cap. Two 20S proteasome subunits (α 4 and β 2) and ten 19S cap subunits are absent, and must be imported like all 11S activator subunits.

Elements of the nuclear pore-complex export/import machinery, the key nucleolar structural protein fibrillarin (Nop1), and elaborate RNA processing machinery confirm that the nucleomorph is a miniaturized nucleus. The surmise¹³ that nucleomorph envelope pores are genuine nuclear pores is confirmed by well-conserved importin genes (*impA*; *imb1*) and a protein interacting with nuclear pores and needed for transport (CRM). Neither transcription nor RNA processing was radically simplified by symbiogenesis. All three RNA polymerases and several transcription factors are nucleomorph encoded, including TfIID for promoter recognition, TfIIB for binding polymerase II, and two components (Rad3; Rad25) of TfIIH. Nucleomorph messengers are probably capped and polyadenylated, for mRNA-capping enzyme (Mce), cap-binding protein (Cbp), and poly(A)-binding protein (Pab) genes exist. Many spliceosomal splicing components (including U6 snRNA and snRNPE, as in chlorarachneans¹²) and some for tRNA intron removal are present, as are 5 RNAs and 17 proteins of the nucleolar snoRNP machinery for processing rRNA (cleavage, methylation and pseudouridylation). The gene for pseudouridylate synthase is of the eukaryotic type (cbf5) with ancillary centromere-microtubule binding properties, but lacks several normally conserved features. The thesis that snoRNAs are ancestral for all cells but were lost in eubacteria by 'streamlining'27 is rendered implausible by their persistence in this ultra-streamlined genome.

Thirty-seven genes encode large subunit periplastid ribosomal proteins and 28 small subunit ones, so about 14 are imported or coded by unidentified ORFs. The nucleomorph encodes 37 tRNAs (12 with introns; Table 1) and standard protein synthesis initiation and elongation factors, but only one amino-acyl-tRNA synthetase (seryl). As we found no gene for glutamyl-tRNA, it must be imported unless its codons are recognized by another tRNA with 'super-wobble'.

Cryptomonad and chlorarachnean nucleomorphs are natural experiments in genome miniaturization and cell simplification that can test basic ideas about genome and cell functions. If domain organization, folding, and mitotic segregation of nucleomorph chromosomes are indeed exceptionally simplified, they could become important models for understanding the more complex chromosomes of typical nuclei.

Methods

Nucleomorph DNA

As nucleomorph DNA is only about 0.1% of cellular DNA, we could not obtain pure enough samples or sufficient amounts of it for a single mechanically fragmented library. We prepared several libraries slightly contaminated by DNA from the other three genomes: random libraries from the nucleomorph DNA band on CsCl density gradients²⁸ (with major mitochondrial DNA contamination) and others enriched in one individual nucleomorph chromosome extracted from bands excised from pulsed field gels⁹. DNA was partially digested by *Sau3*AI and cloned into phage λ as described¹¹, or completely digested with one of *Eco*RI, *Hind*III, *Pst*I, *Bg*III, *XbaI*, *ClaI*, *SpeI*, *BcI*I or *Bam*HI and cloned into plasmids (pUC18).

Sequencing and gene identification

Sequencing, editing and contig assembly were done as described¹¹. Clones were sequenced fully on both strands by primer walking, and gaps between contigs filled by polymerase chain reaction of genomic DNA. Preliminary gene identification and analysis used MAGPIE²⁹. tRNA and snoRNA genes were detected using the search tools tRNAscan (http://www.genetics.wustl.edu/eddy/ tRNAscan-SE) and snoscan³⁰. Further gene analyses used a Perl script to identify all possible ORFs and search them against GenBank (BLAST results available at http:// reith.imb.nrc.ca/nucleomorph.html). Spliceosomal introns were first found manually, and then using a Perl script that searched for conserved splice junctions and extension of the ORF by removing the putative intron.

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Nonlinear effects of large-scale climatic variability on wild and domestic herbivores

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Large-scale climatic fluctuations, such as the North Atlantic Oscillation (NAO)^{1,2}, have been shown to affect many ecological processes³⁻⁶. Such effects have been typically assumed to be linear. Only one study has reported a nonlinear relation⁷; however, that nonlinear relation was monotonic (that is, no reversal). Here we show that there is a strong nonlinear and non-monotonic (that is, reversed) effect of the NAO on body weight during the subsequent autumn for 23,838 individual wild red deer (Cervus elaphus) and 139,485 individual domestic sheep (Ovis aries) sampled over several decades on the west coast of Norway. These relationships are, at least in part, explained by comparable nonlinear and nonmonotonic relations between the NAO and local climatic variables (temperature, precipitation and snow depth). The similar patterns observed for red deer and sheep, the latter of which live indoors during winter and so experience a stable energy supply in winter, suggest that the (winter) climatic variability (for which the index is a proxy) must influence the summer foraging conditions directly or indirectly.

Local weather patterns⁸ and large-scale climatic variability^{6,9–11} affect demography and population dynamics of temperate ungulates. The NAO¹ is a large-scale fluctuation in atmospheric mass between the subtropical North Atlantic region (centred on the Azores) and the subpolar North Atlantic region (centred on Iceland)¹². The NAO is positively correlated with temperature and precipitation at the west coast of Norway during winter¹. Because temperatures in this region often are around 0 °C during winter, precipitation (as rain) is positively correlated and snow depth is negatively correlated with the NAO at low elevation (below 400 m), and precipitation (as snow) is positively correlated with the NAO at high elevation¹¹. Similar differential impacts of global climatic warming at low-altitude and high-altitude habitats have been reported from the Rocky Mountains¹³.

Using wild red deer (*Cervus elaphus*) and domestic sheep (*Ovis aries*) lambs as examples of wild and domestic herbivores, we have explored comparatively the effect of the NAO (being a proxy for the interannual variation in winter temperature and precipitation along the west coast of Norway^{1,11}) on individual performance (measured as body weight). As domestic sheep in Norway are free-ranging only from May/June to September/October, we can separate, through a comparison of the two species, the direct effects (costs of thermo-regulation and movement in snow) and indirect effects (delayed snow melt can affect foraging conditions during summer) of winter climate.

In our study region, sheep and red deer in Norway are sympatric and have very similar diet composition and habitat use¹⁴. During winter, however, sheep are fed indoors, and as a result experience a stable interannual energy supply during that season. Thus, any relation between growth of lambs (during summer) and winter